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International application number: PCT/US04/043969

International filing date: 29 December 2004 (29.12.2004)

Document type: Certified copy of priority document

Document details: Country/Office: US Number: 60/590.987

Filing date: 26 July 2004 (26.07.2004)

Date of receipt at the International Bureau: 09 February 2005 (09.02,2005)

Remark: Priority document submitted or transmitted to the International Bureau in

compliance with Rule 17.1(a) or (b)





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APPLICATION NUMBER: 60/590,987 FILING DATE: July 26, 2004 RELATED PCT APPLICATION NUMBER: PCT/US04/43969

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16805 U.S. PTO PROVISIONAL APPLICATION COVER SHEET This is a request for filing a PROVISIONAL APPLICATION Under 37 CFR 1.53 (b)(2). Type a plus sign (+) 072604 Attorney Docket No. 624.P inside this box ----> INVENTOR(s)/APPLICANT(s) MIDDLE RESIDENCE (CITY AND EITHER FIRST NAME LAST NAME INITIAL STATE OR FOREIGN COUNTRY) Wang Jianying 770 Crane Avenue, Foster City, California 94404 TITLE OF THE INVENTION (280 characters max) HPV INHIBITORS CORRESPONDENCE ADDRESS James J. Wong Gilead Sciences, Inc. 333 Lakeside Drive Foster City STATE California ZIP CODE 94404 COUNTRY U.S.A. ENCLOSED APPLICATION PARTS (check all that apply) х Specification Number of pages 19 Small Entity Statement Drawing(s) Number of sheets Other (specify) METHOD OF PAYMENT (check one) The Commissioner is hereby authorized to charge filing fees (as well as any additional Provisional Filing fees which may be required by this paper) and credit Deposit Account Number 07-1250. Fee Amount (\$) \$ 160.00

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

In re application of: Jianying Wang

For:

HPV Inhibitors

Commissioner for Patents

P.O. Box 1450

Alexandria, VA 22313-1450

PROVISIONAL APPLICATION COVER SHEET (37 C.F.R. § 1.51 (2) (i))

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HPV INHIBITORS

Most Prevalent Serious HPV-Mediated Diseases (US and EU)

- Anal dysplasia in HIV patients
- low & high-grade dysplasias ~ 575,000
- Cervical dysplasia
- high-grade lesions $\sim 470,000$
- low-grade ~ 12MM
- Genital warts
- ~2.3MM
- 280,000 patients/yr receive topical pharmaceutical therapies
 - Anal dysplasia in homosexual males
- low & high-grade dysplasias ~ 700,000

Why is Gilead Interested in HPV

- Series of nucleotides have in vitro activity in HPV+ cell lines
- Cidofovir and analogs effective in animal models

SiHa xenograft and CRPV models

- Cidofovir has shown some efficacy in HPV associated numan diseases
- anogenital warts, cervical intraepithelial neoplasia, respiratory papillomatosis
- HPV-associated diseases are an unmet medical need
- Some indications, particularly anal dysplasia in HIV patients, are a good corporate fit

Human Papillomaviruses

- Small, non-enveloped, DNA virus
- Dependent on host cell for replication
- Epithelial tropism
- site of occurrence (cutaneous, mucosal)
- Species specific with > 100 human subtypes
 - low and high risk subtypes

HPV Genotypes

- High risk:
- HPV-16, 18, 31, 33, 35, 45
- potential to induce malignant proliferation
- HPV-16,18 responsible for 50-80% of dysplasias untreated dysplasias may develop into cancer
- Low risk:
- HPV-6, 11
- responsible for nearly 90% of genital warts

Target Product Profile Overview

- **HPV Activity**
- activity against HPV-16,18; ideally activity against HPV-6,11
- Selectivity
- good selectivity between HPV-infected and non-infected tissue
- Safety
- minimally irritating to mucosal tissue
- non-mutagenic
- ideally, non-teratogenic
- Dosing
- once-daily dosing acceptable for anal and cervical dysplasias and for genital warts
- **Formulation**
- topical gel/foam/cream

Goals for Program

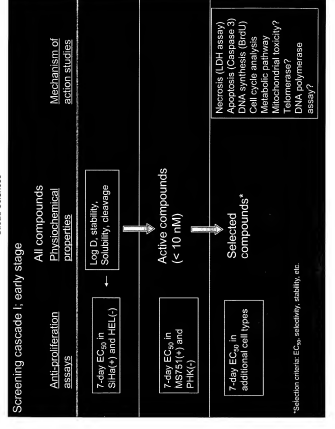
- Topical prodrugs
- improve potency
- allow for skin penetration
- reduce toxicity

Selectivity in vitro

- EC₅₀ "normal" cell line/EC₅₀ HPV+ cell line
- Topical efficacy in representative animal models
- Compound with minimal irritation and minimal/no genotoxicity

Key Challenges

- Selectivity index is low in vitro
- Mechanism of action is not understood
- The parent molecules are potentially toxic
- renal toxicity, mutagenicity, carcinogenicity, local irritation



human xenograft model CRPV model, HPV-6 SiHa xenograft model in mice Animal models Regaj [NIAID] Î In vitro skin permeation Physical and biological Prototype topical Formulation half life in topical formulation formulation Equal or more potent in rabbits than Cidofovir Screening cascade II; late stage Genotoxicity assay Local irritation Toxicology Local irritation Suitable for topical formulation Minimal skin irritation Minimal genotoxicity Minimal genotoxicity compounds from Minimal irritation Skin penetration in vitro studies Selected

cPrPMEDAP: Scaffold for Prodrug Design

- Anti-proliferative activity against HPV(+) cell lines (EC₅₀ at sub-µM range)
- Selectivity when compared to HPV(-) cell lines or primary human keratinocytes

Disadvantages of Other Parent Scaffolds

- Cidofovir (HPMPC) has moderate activity but less selective
- PMEA is not active
- PMEDAP and its other analogs are less active.
- PMEG has similar activity; synthetically more challenging

Improved Potency of cPrPMEDAP Prodrugs

#S9	R	R2	EC ₅₀ (nM) in HPV16+ SiHa cell line
8369	ЮН	Ю	284
17429	O-iPr	O-iPr	2267
327353	Ala-Pr	Ala-Pr	2.5
327238	Ala-iPr	Àla-iPr	1.3
327319	Aba-Et	Aba-Et	3.2
327261	Aba-Bu	Aba-Bu	0.20
327352	OPh	Ala-Pr	0.50
327383	OPh	Aba-Bu	0.13
56884	OPh	Phe-Et	09:0

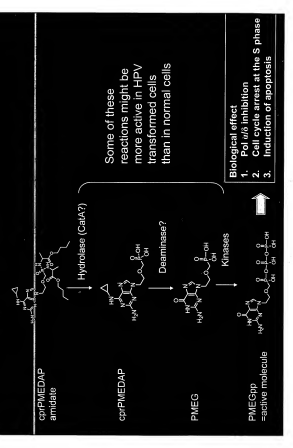
Selection Criteria for Initial in vivo Screens

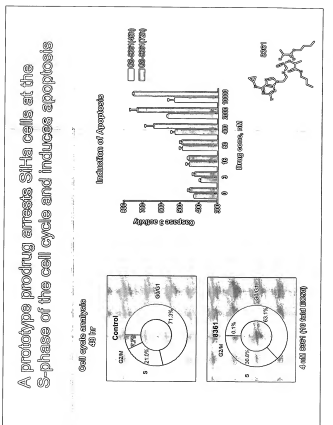
- Key criteria
- Selectivity HPV+/HPV- cell lines
- Log D between 1.5 and 2.5
- Stability in formulation vehicle
- l Other
- Potency in HPV+ cell lines
 - Solubility
- Cleavage by CAT A

CPrPMEDAP Prodrugs with Good Selectivity

#S9	Strı	Structure	Selec	Selectivity
	R1	R2	HEL/SiHa	PHK/SiHa
8369	Ю	HO	17	13
327353	Ala-Pr	Ala-Pr	210	31
327238	Ala-iPr	Ala-iPr	559	75
327319	Aba-Et	Aba-Et	135	12
327261	Aba-Bu	Aba-Bu	115	4
327352	OPh	Ala-Pr	164	10
327383	OPh	Aba-Bu	92	22
56884	OPh	Phe-Et	72	58
Podofilox	Active ingredi	Active ingredient of condylox	<0.9	0.1
8358	PMEG b	PMEG bis Ala-Bu	11	1.8
AraC	C-analog [C-analog DNA pol inh	0.11	0.57
Cladribine	A-analog [A-analog DNA pol inh	pu	1

Mechanism of Selectivity (hypothesis)





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Mechanism of Selectivity (experimental plan)

- Compare the rate of metabolic conversion in cells to identify the rate-limiting reaction
- SiHa (HPV16, sensitive to cprPMEDAP amidates)
- CaSki (HPV16, resistant)
- Primary keratinocytes (HPV neg, somewhat resistant)
- Primary fibroblasts (HPV neg, resistant)

PHK-SiHe Reift Co-cultures (effer 10 days of differentation)



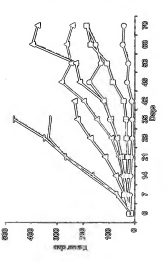


CPrPWEDAP 0.05 µg/mL

CPrPWEDAP 0.5 µg/ml

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Sitta Tumor Model - Athymie Mies Monoamidates ((Ale-îPr))



--- 77384 0.8mg/ml

74 J

3 weeks treatment, 5 days/week

100 pd, intratumoral

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